Collision Detection and Administration Methods for Many Particles with Different Sizes

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Abstract

This paper deals with the calculation of the motion and the administration of the contacts for systems with many colliding bodies of round shape and possibly large size-differences. Both two dimensional (2D) and three dimensional (3D) cases are investigated, while the efficiency of the employed algorithms is compared. For the integration of the equations of motion, standard methods from Molecular Dynamics (MD) and Discrete Element Methods (DEM) are used and, to reduce the effort that is typically used for collision detection, some sophisticated administration algorithms for the neighborhood search are implemented. Especially for large systems with many particles with a wide, polydisperse size distribution, this is a challenge. In the following three methods, the Verlet-Neighbor List (VL), the Linked Cell (LC) method, and the Linked Linear List (LLL), are discussed and compared for 2D and 3D. Only LLL performs well for strongly different particle sizes.

Key words: contact detection; neighborhood search; molecular dynamics (MD); discrete element method (DEM); polydisperse size-distribution

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1 Introduction

In order to determine the dynamical behavior of systems consisting of many objects, particles or atoms, several fully developed approaches exist. The main differences are the assumptions about the particle shapes and their behavior on collisions. Here we examine spherical particles, that could be treated as (i) perfectly rigid objects, as (ii) non-deformable objects with (small) overlaps at the contacts, or (iii) as deformable bodies with a peculiar contact dynamics. Systems consisting of bodies with negligible deformations, i.e., case (i), can be described by means of the so-called multibody system method (MBS) [12, 17], and mass point systems may be regarded as a special case of the MBS. Using a molecular dynamics (MD) [1] also called the discrete element method (DEM), i.e., case (ii), fake-deformable bodies can be represented by a collection of non-deformable particles connected by springs [14]. The difference between MD and DEM are the interaction forces between the particles, otherwise both methods are similar in spirit. For completeness, we remark that the so-called event-driven (ED) molecular dynamics [4, 8, 15] also assumes hard (rigid) spheres and thus can be seen as bridging the gap between MD/DEM and MBS for some special and simple situations. For more advanced studies of flexible, deformable bodies, i.e., case (iii), usually the Finite-Element-Method (FEM) or the Boundary-Element-Method (BEM) are used, see [2, 13]. Each of these methods has its own advantages and disadvantages. While the MBS is in general characterized by relatively short computation times due to a small number of degrees of freedom, deformations cannot be handled. On the other hand, systems investigated using FEM have a large number of degrees of freedom that yield a rather extensive number of equations of motion, but deformations are properly taken into account. The MD/DEM methods can be seen as a compromise in so far that the number of degrees of freedom is kept small by assuming overlap dependent contact force laws that rely on certain assumptions and do not take the eigen-modes of a particle into account. An expansion of the MBS method for elastic bodies is e.g. presented in [9].

Hybrid MBS/FEM contact calculations are presented in [2], where colliding bodies are examined by the FEM approach in order to incorporate deformations while all the other bodies of the system are regarded as rigid. This approach combining FEM and MBS makes use of the advantages of both methods. However, these advanced (hybrid) approaches share the drawback that the number of contacting particles is quite limited due to the detailed modeling involved for each particle.

Alternatively, very efficient methods were developed for molecular dynamics and discrete particle systems, i.e., granular matter [3, 15, 20]. Molecular (gases, fluids, solids) and particulate systems, charged or neutral, can be investigated [1, 7, 15]. The motion and the contacts of many thousands of particles, involving body forces like gravity and interactions with the boundaries of the system, can be modeled. The formulation of the interaction forces between
the different bodies is based on models as simple as possible, in order to keep the calculation time feasible. Usually, very small penetration of otherwise non-deformable particles are accepted, compare [1, 5, 16, 20], and are used as the basis of the force calculation. One of the essential assumptions, which allows for efficient algorithms for granular systems, is that the particles interact only when in contact.

For a given pair of particles, a normal contact force acts in the direction opposite to the penetration—along the center-to-center line, and is usually modeled as a spring dashpot element that combines elastic and viscous response on the contact level. The spring force is proportional to the penetration of the particles, see [5, 11], corresponding to a penalty force. The simplest force model is linear, but also various non-linear models are available [5]. Apart from the repulsive, and viscous contact forces, also attractive forces may occur due to adhesion/cohesion, or electro-static interactions. The latter are typically long-ranged and are not subject of this study, where we focus on short range, contact interactions only. These also could involve plastic deformation and friction as alternative sources for dissipation; we refer to Ref. [6] for more details and restrict ourselves to the most simple, linear contact forces in the following.

For a system consisting of \( n \) particles with arbitrary interactions, the required calculation operations for the force computation will be of the order \( O(n^2) \), causing huge computational effort. However, for the systems with short-range contact forces, only particles in their respective neighborhood can interact, so that a tremendous reduction of computational effort down to the order \( O(n) \) can be achieved [1].

In this study, we consider \( n \) spherical bodies with radii \( r_i \), consecutively numbered from \( i = 1, \ldots, n \), where the radii can be different. The used contact force model is based on linear spring dashpot elements, so that the equations of motion are

\[
m_i \ddot{r}_i = \sum_j \left[ k \delta_{ij} + d \dot{\delta}_{ij} \right],
\]

with the masses of the bodies \( m_i \), the positions \( r_i \), and the forces \( f_{ij} \) acting between \( i \) and \( j \), while the sum runs over all particles \( j \) in the neighborhood of particle \( i \). Here, \( k \) is a spring constant, while \( d \) is the damping coefficient for the dissipative force. The overlap between two particles \( i \) and \( j \) is

\[
\delta_{ij} = r_i + r_j - (r_i - r_j) \cdot n_{ij},
\]

with the normal vector between both particles, \( n_{ij} \), parallel to the line connecting their centers. These contact forces are applied only for the case \( \delta_{ij} > 0 \). Frictional forces as well as adhesive contact forces are not considered here. The equations of motion are solved by means of the Verlet integrator, which is an
explicit integrator. The new positions of the particles are computed based on the information about the actual positions and the positions during the previous time step, that means without knowledge of the velocities of the particles. In section 2, different approaches to save computational effort as mentioned above are introduced and discussed. In section 3 these methods are compared using several test examples and finally, in section 4, the results are summarized and discussed with respect to possible applications.

2 Neighbor Search Methods

The following three methods presented can be used in order to find neighboring bodies of a particle efficiently. Two of these methods, VL and LC, identify the neighboring particles of a body by regarding special regions of the system and considering all particles within the same region as neighbors. As both methods operate with similar ideas, they are also sometimes used simultaneously [16]. The third method, LLL, is based on a different approach. Around each body a bounding box is placed which is proportional to the particle size; every body whose bounding box is colliding with the bounding box of another particle is considered to be a neighbor of this particle and thus a potential contact partner [18].

For each method the neighboring particles are stored in a neighbor data-structure (NDS) after some pre-sorting. The collision detection needs to be done only for neighboring and potentially colliding bodies. Hence, the necessary calculation operations for collision detection can be reduced down to an order proportional to the number of particles in the system, i.e. \( O(n) \), because the update of the NDS can be optimized. In the following we will examine whether this expected performance also holds for LLL – but first the three methods are introduced.

2.1 Verlet-Neighbor List

The first method presented, the Verlet-Neighbor List (VL), is quite similar to Verlet’s originally proposed method. As shown in Fig. 1 an imaginary sphere is drawn around each particle of the system, e.g., with a radius of five times the maximum radius of the particles, compare [1, 10]. Particles within these enclosing spheres are considered as neighbors of the particular body in the middle of the sphere. The optimal extension of the zone around the bodies depends on the velocity of the particles and on the density of the whole system. For each particle a list is generated, where all neighboring bodies are stored [21]. In order to compile these neighbor lists, for each body all particles of higher numbers than the body itself have to be tested, whether they lie inside the test
sphere or not. Particles of lower numbers are not tested since no pair needs to be checked twice.

Creating these neighbor lists therefore requires \(n(n-1)/2\) calculation steps, which means the number of necessary operations is still of order \(O(n^2)\). However, the lists do not have to be updated for every time step. The update frequency depends on the density of the system, the velocity of the particles, and on the size of the spheres. For relatively dense systems, an update of only each 100th time step can be enough. Like the value for the radius of the spheres around the particles, also the update frequency is a parameter that can be tuned and, in principle, has to be checked for every new simulation. Both parameters are interdependent, because the value for the radius is inversely related to the rate at which the list must be rebuilt, see [16]. The smaller the zone around the particles, the more often the reconstruction of the lists needs to be done. The larger it is, the more particles belong to the neighborhood requiring more contact calculation time.

The real collision detection and force calculation, that requires additional effort, especially for more advance interaction models, now only has to be done for the particle pairs which are stored in the lists. That leads for the example of Fig. 1 to just four pairs (and force calculations) instead of the originally required ten.

2.2 **Linked Cell Method**

An alternative approach that is often used to indentify the neighbors of a body is the Linked Cell (LC) method, where the system is divided into a regular lattice of, e.g., for cubic systems, \(m \times m \times m\) cells (3D) [1]. Non-cubic systems can also be observed and the cell shape can be chosen as to fit the system with more or less cubical cells. The optimal size of the cells, as before the size of the Verlet spheres in the VL method, depends on the velocity of the particles and on the density of the system, but the cell sizes all need to exceed at least the size of a particle in length. The major difference between LC and VL is, that for LC the cells are not sticked to particles and thus are
not moving along with the particles. The size of the cells can be selected such that one cell contains about three particles, see Fig. 2. Then, if particles are temporarily assigned to special cells on the basis of their current positions, it is obvious that interactions are only possible between those in the same or in directly adjacent cells [16]. This means for a 2D system, that only particles within nine different cells for 2D and 27 cells for 3D may contain neighbors. Again, as in the previous description for VL, particle pairs are only checked once. Thus, not all cells have to be tested, but only the center cell plus half of the neighboring cells. That means one plus four (2D), Fig. 3, or one plus thirteen cells (3D), Fig. 4. Cast into a formula, we have \((3^d + 1)/2\) cells to examine, where \(d = 2\) for a 2D system and \(d = 3\) for a 3D system.

For the 2D system illustrated in Fig. 2 the NDS for body 1 contains the
particles from 2 up to 15. For particle 2 all particles of the neighboring cells except number 1 are neighbors and for particle 6 only particles of the four neighboring cells are neighbors. This is due to the fact that also within a cell the same strategy is applied in order to avoid multiple checks of the same particle pair.

In such a $d$-dimensional system, on average, one has $n_c = n/m^d$ particles in each cell, with $n$, the number of bodies in the whole system and $m$, the number of cells in each direction. Therefore, as an estimate, on average, only

$$p \approx \left[ (3^d - 1) \frac{n_c^2}{2} + \frac{n_c(n_c - 1)}{2} \right] m^d = \left[ 3^d \frac{n_c^2}{2} - \frac{n_c}{2} \right] m^d$$

(3)

distances need to be examined (for the majority of cells that are not situated along the edge of the system). The first term takes neighbors outside into account, while the second those inside the cell of the particle of interest. The factors $1/2$ are due to the fact that every pair shall be checked only once. Since the bracket in Eq. (3) equals the approximate number of necessary operations for one cell, it has to be multiplied by the number of cells in the system, $m^d$.

Another estimation for the average number of pair checks was given in [1] as

$$p \approx 4.5nn_c,$$

(4)

for 2D, which is almost identical to Eq. (3) for large $n_c$. For each particle, in half of the nine cells approximately $n_c$ particles will be neighbors. This contrasts with

$$p = \frac{1}{2} n(n - 1),$$

(5)

which is of order $O(n^2)$ if no neighborhood search method is used. Clearly, for systems with $m \leq 3$, there is no benefit in using any NDS, but usually the systems observed possess much more cells, if the number of particles is high enough to justify a method like the described ones.

2.3 Linked Linear List

The third possibility described in this comparison, also a very efficient method used to keep track of neighbors for large systems, is the Linked Linear List (LLL) [18]. This method is quite different to the other approaches. In a first step, bounding boxes are laid around each particle, Fig. 5, that are sized in such a way, that each particle fits exactly in its box. The edges of each bounding box are aligned parallel to the system axes.
**Linear List Generation**

In a next step the bounding boxes are projected separately onto the system axes. Such a projection onto the x-axis for the situation in Fig. 5 is shown in Fig. 6. In the following, only the order of the beginnings ‘b’ and endings ‘e’ of the projections of the bounding boxes along the axes is of interest. For this reason the sequences are stored in lists.

Time $t$

\[
\begin{array}{cccccc}
1 & 2 & 3 & 4 \\
\text{b}_1 & \text{e}_1 & \text{b}_2 & \text{b}_3 & \text{e}_3 & \text{b}_4 & \text{e}_2 & \text{e}_4 \\
\end{array}
\]

Time $t+\Delta t$

\[
\begin{array}{cccccc}
1 & 2 & 3 & 4 \\
\text{b}_1 & \text{e}_1 & \text{b}_2 & \text{b}_3 & \text{b}_4 & \text{e}_3 & \text{e}_2 & \text{e}_4 \\
\end{array}
\]

Fig. 6. Particles projected on the x-axis for two different times, [18].

For a 3D system, three different projections are necessary and, therefore, three lists will be generated. Each of them has a length which corresponds to twice the number of particles in the system. If there is the beginning, ending, or both, of another particle in between the beginning and ending of a particular body, then there will be an overlap of the projections of the bounding boxes of both particles along this axis. A collision of two bounding boxes exists for an overlap of these projections along each axis.
Update of the Linear Lists

Checking whether there is some part of a projection in between the beginning and ending of another projection for each particle along each axis still takes a lot of time. But, although these lists have to be updated for each time step, the necessary calculation times can be reduced to an amount proportional to the total number of particles in the system, as there has to be done only an update of the old list for each new time step. That corresponds to sorting an already nearly sorted list. This update can simply be done by going through the lists sequentially and checking for any new changes in the order. The occurring changes are usually permutations only, compare e.g. Fig. 6, where $e_3$ and $b_4$ have been changed. If the order of the beginnings and endings does not have to be changed, the collision status of the particles also will remain unchanged. While seeking for new colliding bounding boxes by looking for permutations in the lists, four different cases have to be discerned, compare [18].

1. Two beginnings are changed, which means the bounding boxes have been overlapping and continue to overlap.
2. Two endings are changed, which also means the bounding boxes have been overlapping and continue to overlap.
3. A beginning and a proximate ending are changed, which means a so far occurring overlap has to be removed.
4. An ending and a following beginning of another particle are exchanged, which means a previously non-existing overlap has to be taken into account.

For the first two cases, except for the exchange, nothing has to be done in the lists, as the collision status between any particle will not change. If a collision along an axis has to be removed, or if there is a new collision between two particles along an axis, the collision information along the other axes is essential. One has to know whether there is a new or old collision along all axes and, therefore, between the bounding boxes or, whether there is no overlap any more between two so far colliding bounding boxes in at least one direction. For this reason, a second and a third column (for the 2D case) are added to the lists, that store the information of the positions of beginnings and endings along the y-axis, see Fig. 7, respectively. In each row, the positions are stored, of beginnings (column two) and endings (column three) of the particle of the first column. For a 3D system also a fourth and a fifth column with the position information of beginnings and endings along the z-axis have to be added. For our 2D example going through the list along the y-axis, see Fig. 7, leads to the potential collision between particles $(3/4)$, and $(1/2)$. As the location of particle 3 along the x-axis is from position three to five, whereas the beginning of particle 4 has the position four, there is also an overlap of bounding boxes 3 and 4 along the x-axis and, therefore, a real collision of the bounding boxes of particles 3 and 4.
Fig. 7. Lists containing also the position information along the other axes.

Particles 3 and 4 are now considered to be neighbors, that have to be checked for collision. Therefore, a linked list is created in the form of a sparse matrix where the colliding bounding boxes are stored. Collision pair (3/4) is stored at position 3, 4 of the matrix, see Fig. 8, that shows particle pair storage for an arbitrary configuration.

Fig. 8. Storage of colliding bounding box pairs, if e.g. boxes (1/2), (1/3) and (3/4) collide.

Figure 9 shows an example of a 2D system and Fig. 10 the matrix structure that is received for this system. As the particle numbers are given sequentially, the entries in the matrix for the left part of Fig. 10 that are right next to the diagonal, show that only neighboring bodies have colliding bounding boxes. This is only true for the special numbering, that we use at the beginning of our simulations, but the matrix is for all times quite sparse, cp. right part of Fig. 10.

As collisions between the bounding boxes of two particles are only treated once, in Fig. 10 an upper triangular matrix is shown, where the diagonal of the matrix must also be empty. Due to the very small number of entries in the matrix, this cannot be seen very clearly in this illustration, but it is clear from Fig. 8. All rows and all columns are linked, compare Fig. 8. Therefore, it is
Fig. 9. System consisting of 300 particles for two different points of time. The initial situation is shown on the left hand side, whereas the right side shows the situation after a simulation time of 2 sec, calculated with gravity, but without dissipation and friction.

Fig. 10. Matrix structure for two different times, for the 300 particle system, where the dots are the potential collisions for the situations in Fig. 9.

now possible to check only these neighboring particles in a following collision fine test, whose bounding boxes collide.

2.4 Discussion

The two techniques described in sec. 2.1 and sec. 2.2 have in common that both identify neighboring particles by considering bodies inside certain zones as neighbors. For both methods these zones have to be at least somewhat larger than the particles themselves. From this, two problems can arise. Firstly, if the particles within the system are polydisperse, that means their sizes differ, then the size of the grid (LC) or circles (VL) has to conform to the largest particle existing in the system. Hence, for highly polydisperse mixtures, the smaller
particles may increase the number \( n_c \) of particles within one cell, which might even, in the worst case, be close to \( n \), see [18].

As the neighborhood zones around a particle are larger than the particles, the NDS does not have to be updated in each time step. The size of the zones and the necessary update frequency are interdependent and not quite easy to guess. Therefore, another problem of both methods is the ascertainment of optimal values for both, the update frequency for the lists and the size of the zones. If the cells are chosen smaller than the particle size, a successful contact detection can be impossible in the way described above. Besides that, if the cells are either very small or very large, the contact detection is inefficient because updates are too frequently necessary or too many particles are in the neighborhood, respectively. Therefore, the choice of these values is very important, and it may take a lot of personal time getting experience with the investigated system.

3 Comparisons and Results

The three techniques introduced shall in the following be compared with respect to the simulation times. In order to keep the influence of the different computers, of the different programming languages and compilers, as well as the influence resulting from the different programming styles as small as possible, for the comparisons different test series are used. The goal of these series is, to keep some system traits as constant as possible and to change only some well defined influencing factors.

The program for the LC was programmed in C++ while the VL and LLL were programmed in C. For this reason and since the program runs were performed on different Linux PC’s we normalized the results in such a way that for the smallest investigated number of particles for each system (where the influence of the used method is mainly negligible) contact scaling factors have been computed. Therefore, all curves in the following figures have the same starting point.

3.1 Comparison of a Planar Polydisperse Example

In a first 2D test series, examples with different numbers of particles are studied. The system is not monodisperse, i.e. the diameters of all particles are not equal, but the difference in the particle sizes are equal for each example of the series. The sizes were randomly drawn from a homogeneous distribution in the interval \([R_0(1-w_0), R_0(1+w_0)]\) with mean radius \( R_0 = 10^{-3} \text{ m} \) and width \( w_0 = 0.5 \). Other system parameters are the stiffness for the calculation of the penalty force \( k = 10^7 \text{ N/m} \), the dissipative constant \( d = 0 \text{ Ns/m} \), the
density of the particles $\rho = 7000 \text{ kg/m}^2$, and the time step for the integration of $dt = 10^{-7}$s. It was tried to keep the density of the system as constant as possible, while simultaneously the number of particles was duplicated stepwise and the system size was enlarged accordingly, see Fig. 11.

![Fig. 11. Example of a dense system with 7800 particles. The lines show the arrangement of the particles for all six system sizes.](image)

Keeping the system density for each example exactly constant was not possible, since the container size divided by the linked cell size always has to be a natural number and the linked cell size for the whole series was kept equal $l_c = 7.0 \times 10^{-3}$ m. The first series consists of

244,
488,
975,
1950,
3900, and
7800 particles, respectively.

The results, received for the computation times with respect to the increasing number of particles are shown in Fig. 12. From this picture it is quite clear, that the curve for the Verlet method is the steepest, whereas both other curves have a very similar behavior. All three curves show the most increasing behavior between 1950 and 3900 particles. However, reason for this behavior are slight changes in the density of the system, which varies from $\nu_{cont} \approx 0.670$ to $\nu_{cont} \approx 0.678$. In Fig. 13 the curves for LLL and LC are pictured enlarged again. Supplementary there is a straight line added to the graphs, that is parallel to both curves below 1950 particles. The picture shows, that the gradient of the curve for LLL between 3900 and 7800 particles is very similar to the gradient below 1950 particles and that the gradient for LC above 3900 particles is even lower than it was for the smaller systems. Hence, from the gradient of the straight line it is clear, that the gradient of both curves is approximately

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equal two, and that an $O(n)$ runtime behavior is obtained.

3.2 A Spatial Monodisperse Example for Increasing System Size

A series of 3D systems with an increasing number of particles of equal size, $R = 5.0 \times 10^{-4}$ m, and a proportionally growing space around the particles is investigated in this subsection. For the stiffness of the particles it was chosen $k = 10^5$ N/m, the damping coefficient again was neglected, the density of the particles was $\rho = 10^{10}$ kg/m$^3$, and the time step for the integration was chosen
as $dt = 10^{-5}$ s. Here, systems of

512,
2197,
5832,
12167,
21952,
35937, and
59320 particles

are investigated, where again the linked cell size was kept equal $l_c = 2.0 \times 10^{-3}$ m for each system. The behavior of the computation time with respect to the number of particles within the particular system is shown in Fig. 14 as log-log plot.

![Graph](image)

Fig. 14. Comparison results for an increasing 3D example.

The more the number of particles rises, (and with it the number of contacts during the simulation) the more the performance for VL escalates in comparison to LLL and LC. Here LC has the best performance. Clearly it can be seen, that the gradient is lower than for LLL. Due to clearness of the plot the approximating curves are not plotted in Fig. 14. Nevertheless, a roughly similar curve for the behavior of VL is $y \approx (0.027 \, x)^{1.8}$, the polynomial for LLL is $y \approx (0.12 \, x)^{1.36}$ and the polynomial for LC is $y \approx (0.2 \, x)^{1.15}$. Therefore, it can be said, that the behavior of LLL and LC both remain close to linear in contrast to the behavior for VL, which behaves almost quadratic.
In the last 3D comparison the system size is kept constant. In contrast to the previous changes, here the number of particles is increased, but no change of either the system or cell-size is undertaken, $l_c = 0.033$ m. Chosen parameters for this system are the stiffness $k = 4.0 \times 10^6$ N/m, damping coefficient $d = 0$ Ns/m, density of the bodies $\rho = 7000$ kg/m$^3$, and the time step for the integration $dt = 4 \times 10^{-7}$ s. The systems can be seen as a series of fracture of some of the particles, where neither the volume enclosed in the system boundaries nor the mass content of the system is changed. That means, that the volume fraction and the density of the system are unchanged, while the number of particles within the system is increased. In the first system 1000 particles are situated, with equal radii $R = 0.01$ m. Approximately half of these particles are now successively fractured in the next systems successively: There are about 500 particles of radius $R$, but approximately eight times 500 particles of radius $R/2$ and thus an eighth of the original particle volume. The systems therefore contain

- 1000 particles (see Fig. 15 on the left),
- 4451 particles, about 4000 smaller bodies of $r = R/2$,
- 14676 particles, about 14000 particles of $r = R/3$,
- 32374 particles, about 31000 particles of $r = R/4$, and
- 65480 particles, about 65000 particles of $r = R/5$ (see Fig. 15 on the right).

![Fig. 15. A monodisperse system and a very polydisperse system with $r = R/5$ of exactly the same volume and density (volume-fraction $\nu = 0.12$).](image)

Here, $R$ is the original radius of all particles of the first system and $r$ the second, smaller one. In Fig. 15 there are no overlaps between the bodies. The bodies that look as if they were overlapping are situated in different depths. For this system the computation time needed per particle is presented in Fig. 16 over the ratio of the radii that is equivalent to the polydispersity of the system.
It can be seen that the gradient of the curve over the polydispersity of the LC calculation is a lot higher than for LLL. Before scaling, so that both curves have the same starting point in the picture, the starting point of LC (for the monodisperse system) was underneath the starting point for LLL, and the curves were intersecting at about $R/r = 2.5$. This means that the LC method has advantages for quite monodisperse systems, while the LLL shows its advantages for polydisperse systems.

### 4 Conclusions

In this paper three different methods have been introduced in order to reduce the calculation times for collision detection for systems consisting of many particles with contact interactions only. The basic idea of these techniques is the fact, that usually there are lots of particles in a system, which cannot be in touch, as they are too far apart. The presented methods save a lot of time by excluding such particles from a detailed and time consuming contact examination and evaluation.

It has been shown, that for rather small systems the traditional VL is a quite good technique, its efficiency being reasonable as it is very easy to implement. For larger systems the LC and LLL methods become more and more efficient compared to the VL. The LC method shows very good performance with a lower increase of computing time for monodisperse systems, as compared to the LLL method. A problem using LC might be the optimal choice of the linked cell size. For too small sizes and for too large sizes the results become inefficient. An optimal cell size has to be found for each different system, since there is no general rule to our knowledge.

As the linked cell size is dependent on the largest particle in the system and as the calculation time is dependent on the linked cell size due to increasing
with increasing cell size, LC becomes very inefficient for wide size distributions. The time spent using LLL behaves much better, as compared to the LC, for polydisperse (partial fracturing) systems of constant volume and density but increasing numbers of particles. Therefore, there is no unique “winner” of our comparison and for computations it is required to know the methods with their strengths and weaknesses in order to choose the most appropriate one.

In future work it might be interesting to use LLL for polydisperse systems with even wider size distributions. Another interesting task will be the contact detection and force calculation of polygonal/polyhedral particles and the neighborhood search for strongly anisotropic particles. Finally, since the collision detection for concave particles is even more complicated and time consuming, the choice of the most appropriate neighborhood search algorithm will remain an issue of interest and requires further research.

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<td>Lists containing also the position information along the other axes.</td>
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<td>8</td>
<td>Storage of colliding bounding box pairs, if e.g. boxes (1/2), (1/3) and (3/4) collide.</td>
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<td>9</td>
<td>System consisting of 300 particles for two different points of time. The initial situation is shown on the left hand side, whereas the right side shows the situation after a simulation time of 2 sec, calculated with gravity, but without dissipation and friction.</td>
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<td>Matrix structure for two different times, for the 300 particle system, where the dots are the potential collisions for the situations in Fig. 9.</td>
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Computation time for a polydisperse system of constant density.